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(54) Title: HIGH PURITY COMPOSITION COMPR	ISING	(7	$7\alpha,17\alpha$)- 17-HYDROXY- 7-METHYI	- 19-NOR-17-PREGN-

(54) Title: HIGH PURITY COMPOSITION COMPRISING (7α,17α)— 17-HYDROXY— 7-METHYL— 19-NOR-17-PREGN-5(10)-EN-20-YN-3-ONE

(57) Abstract

The invention pertains to a process for the preparation of a high purity composition of $(7\alpha,17\alpha)$ – 17-hydroxy– 7-methyl–19-nor–17-pregn– 5(10)–en–20-yn–3-one. The process provides for a composition with less than 0.5 % of $(7\alpha,17\alpha)$ – 17-hydroxy–7-methyl–19-nor–17-pregn– 4-en–20-yn–3-one. This composition can be used as a source for the preparation of stable pharmaceutical base units.

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High purity composition comprising (7α,17α)-17-hydroxy-7-methyl-19-nor-17-pregn-5(10)-en-20-yn-3-one

The invention relates to a high purity composition comprising $(7\alpha,17\alpha)-17$ -hydroxy-7-methyl-19-nor-17-pregn-5(10)-en-20-yn-3-one, a method for the preparation of this compound for use in the pharmaceutical composition as well as a pharmaceutical composition prepared by admixing a pharmaceutically suitable carrier and the high purity composition.

The compound $(7\alpha,17\alpha)$ -17-hydroxy-7-methyl-19-nor-17-pregn-5(10)-en-20-yn-3one (Tibolone) having the structural formula 1:

Formula 1

is known, for example from US 3,340,279 and US Patent 4,701,450. The method described in these patents leads to a compound having combined oestrogenic, progestagenic and androgenic characteristics. This compound is used in medicaments having gonadomimetic, ovulation-inhibiting or immuno-modulating action.

Compositions comprising Tibolone and a pharmaceutically acceptable solid carrier have been described in EP 389 035, which disclosure is incorporated herein by reference. Tablets are available on the market under the name of Livial[®].

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The known tablets can be stable stored very well for, typically, 2 years at ambient temperature. A sufficiently humid atmosphere (e.g. 50 - 70 % relative humidity) makes for a better storage stability than a relatively dry atmosphere (e.g. 45% relative humidity or below that).

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A problem in the preparation of pharmaceutical dosage units is that during the preparation the relative amount of impurities may increase. In particular, the amount of one of the impurities which is already present in the bulk preparation i.e. $(7\alpha,17\alpha)-17$ -hydroxy-7-methyl-19-nor-17-pregn-4-en-20-yn-3-one (Org OM38) tends to increase during the process of making pharmaceutical dosage units. It is furthermore known that the amounts of Org OM38 in compositions comprising Tibolone increase upon storage.

The end of shelf life specification with respect to the amount of Org OM38 formed during storage is 5%. A minimum acceptable shelf life period for these dosage units is 1 year. It is an object of the present invention to improve upon the storage stability i.e. to enhance the shelf-live of the dosage units.

The customary amount of Tibolone in the known dosage unit is 2.5 mg in tablets or capsules of 100 mg, i.e. 2.5%. For the sake of providing therapies better tailored to the individual woman's needs, it is desired to provide dosage units having a lower amount.

However, adaptation of a known formulation by simply including a lower amount of Tibolone further decreases the stability of the dosage unit substantially. E.g., if a 2.5 mg Tibolone dosage unit has a shelf-life of, e.g., 2-3 years at room temperature, the same unit upon lowering the amount of Tibolone to e.g. 0.3 mg can only be kept at 4°C for a period of 6-12 months. Such a lower stability is unacceptable in daily practice. It is a further object of the invention to provide dosage forms having a lower content of Tibolone (which are more prone to stability problems than regular dosage forms) and that can be suitably kept for a prolonged period of time.

One of the possibilities to keep the amount of Org OM-38 below a desired level also after a prolonged storage time is to limit the amount initially present in the bulk preparation. Thus, there is a need to synthesize high purity Tibolone batches with a low contamination content of Org OM-38. It is an object of the present invention to provide for such high purity batches of Tibolone.

During the last step of the synthesis of Tibolone a solution of $(7\alpha,17\alpha)$ -3,3-dimethoxy-17-hydroxy-7-methyl-19-norpregn-5(10)-en-20-yn-3-one in a mixture of pyridine and ethanol is mixed with a solution of oxalic acid in water and the mixture is stirred for 3 hours at approximately 30 °C. The solution is then poured out in a mixture of pyridine and water and the resulting suspension is filtered. The crystals are washed with a mixture of water and pyridine and subsequently, the crystals are dried under

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vacuum at 40 °C to give $(7\alpha,17\alpha)$ -17-hydroxy-7-methyl-19-norpregn-5(10)-en-20-yn-3-one (see also van Vliet et al (1986), Recl.Trav.Chim.Pays-Bas 105, 111-115).

As this compound has a lower stability than the corresponding $(7\alpha,17\alpha)-17$ -hydroxy-7-methyl-19-nor-pregn-4-en-20-yn-3-one there is always formed a small percentage of the latter compound via acid catalyzed isomerisation. Furthermore, this isomerisation takes place at higher temperature and upon long term storage of the crystals

Unexpectedly, it now has been found that the rate of formation of Org OM38 during drying and storage in a specific batch can be decreased if crystals of Tibolone are washed with water and are allowed to age for at least 24 hours in the presence of water. Thus, the Tibolone is left for at least 24 hours under wet conditions. Preferentially the crystals are left under these conditions for a period of at least 3 days. There is no limit to a maximum period but a period of 3-6 days is best suited. The aging temperature preferentially is room temperature.

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Thus according to the procedure of the present invention highly pure Tibolone with a low Org OM38 impurity is obtained by including a delay of several days before drying. The procedure reliably results in batches of Tibolone having a low Org OM38 content. A further advantage is that these batches have an excellent stability. Furthermore, these batches do not form additional amounts of the latter compound upon heating or long term storage.

The crystal formation procedure of the present invention can perfectly well be combined with the last step of the Tibolone synthesis wherein $(7\alpha,17\alpha)$ -3,3-dimethoxy-17-hydroxy-7-methyl-19-norpregn-5(10)-en-20-yn-3-one in a mixture of pyridine and ethanol is mixed with a solution of oxalic acid in water. In general, this reaction proceeds under mild acidic conditions in the presence of an organic solvent and water within a pH range of 5-3, preferentially 3.5-4.5. The acid preferentially is a weak organic acid having a pKa value in the range 1-5 such as citric acid, malonic acid, oxalic acid, dichloroacetic acid and acetic acid, optionally buffered with a base such as pyridine. As organic solvent e.g. ethanol, methanol, acetone, 2-propanol or tetrahydrofuran can be used. The solution is then poured out in water, which is made slightly alkaline by addition e.g. of a low amount of pyridine. After filtering the suspension the crystals are washed with a mixture of water made slightly alkaline by e.g. pyridine. Before drying the crystals are left wet for at least 24 hours.

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Inclusion of the crystal aging step according to the invention results in bulk Tibolone batches with a low Org OM38 content. Routinely, batches are obtained with an Org OM38 content of less than 0.5%. Often even batches with less than 0.25% or even 0.1% of Org OM38 are obtained. Thus high purity compositions with Tibolone having less than 0.5% of Org OM38, preferably 0.25%, more preferably 0.10% of Org OM38 form part of the present invention. The amount of Org OM38 is calculated as the percentage (w/w) of the total amount of the bulk substance including some minor impurities. The amount of Tibolone usually is more than 98%.

The batches of these high purity Tibolone compositions with their low initial Org OM38 content are perfectly well suited to be used as a source for the preparations of pharmaceutical formulations. This guarantees a formulation with a low initial Org OM38 content and improves therefore its storage properties. Pharmaceutical preparations prepared with high purity Tibolone usually result in preparations with less than 1% of Org OM38, often even less than 0.7% of Org OM38 and these preparations are less prone to increase in Org OM38 content during storage.

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As indicated before the amount of Org OM38 in a dosage form also depends upon the concentration of the active substance, the amount of impurity being higher as the amount of Tibolone in the dosage unit decreases. Therefore, using high purity Tibolone as the active substance, dosage units can now been prepared with a lower amount of Tibolone and still having an acceptable shelf life. Thus, the invention also relates to pharmaceutical dosage units, which can be prepared by admixture of a pharmaceutically suitable solid carrier and the high purity composition of the present invention.

A typical known formulation for Tibolone is a 100 mg dosage unit having 2.5 mg of Tibolone contained therein, a relatively small amount (e.g. approximately 1 % by weight) of pharmaceutically acceptable auxiliaries, and a carrier making up the body of the tablet. The carrier typically is composed of 10 % by weight of starch, e.g. potato starch, and 90 % by weight of lactose.

Due to the excellent stability properties of dosage units with a lower amount of active substance than the present commercially available tablets of 2.5 mg active substance, the present invention now makes it also possible to provide for stable dosage units comprising Tibolone in an amount of less than 2.50 mg, preferably 1.25 mg or less, more preferably 0.625 mg or less. At a shelf life of 1.5 years, preferably 2 years these dosage units still comprise less than 5% of OM38 (relative to the amount of Tibolone).

It is another aspect of the present invention to provide dosage units comprising Tibolone in amounts of less than 2.50 mg, preferably 1.25 mg or less, more preferably 10

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0.625 mg or less and comprising at a shelf life of 6 months less than 3 %, preferably 2 % of OM38. The shelf life preferably is extended up to 1 year, preferably 1.5 year, more preferably 2 years.

As used herein shelf life means storage during a specified period under temperature conditions varying from 2-25 °C. Dosage units can be packed e.g. in push-through packs (PTP, blister) and are preferably stored in dark (e.g. enclosed in carton). Alternatively they might also be stored in bottles e.g. high-density polyethylene bottles.

The pharmaceutical dosage units of the present invention will generally take the form of tablets or capsules, but other solid or dry pharmaceutical preparations are included.

Methods for making such dosage units are well known. For example in the standard English language text Gennaro et al., Remington's Pharmaceutical Sciences, (18th ed., Mack Publishing Company, 1990, see especially Part 8: Pharmaceutical Preparations and Their Manufacture), methods of making tablets, capsules and pills and their respective components are described.

Tablets and capsules are prepared of granulates using dry or wet granulation techniques as disclosed in The Theory and Practice of Industrial Pharmacy(Third edition) L. Lachman, H.A.Lieberman and J. L. Kanig (1986) p 1 -99 and 293 - 345.

The aim of granulation is to improve the flowability and compressibility of the powder mixture. Wet granulation forms the granules by binding the powders (a mixture of a diluent and disintegrant) together with an adhesive. The wet granulation technique employs a solution, suspension or slurry containing a binder, which is usually added to the powder mixture; however the binder may be incorporated dry to the powder mix and the liquid may be added by itself. The wet granulation process is performed in mixers/kneaders or fluid bed systems.

Usually an amount of water is incorporated in the basic granulate ranging from 5.5 - 7 %. Preferably the amount of water incorporated is at least 6%.

After granulation the mass is dried to the desired water content using fluid bed dryers, tray dryers, vacuum dryers or other suitable dryers.

To attain a good distribution of the active (Tibolone) over the total mass, the active is premixed with a part of the granulate, sieved using an oscillating sieve, a high speed sieve or other suitable sieving equipment. Next this mixture is mixed with the remaining part of the granulate and a lubricant. This mixture is compressed to tablets, or filled into capsules.

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The following examples are illustrative for the invention and should in no way be interpreted as limiting the scope of the invention.

Examples

Example 1

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A solution of $(7\alpha,17\alpha)$ -3,3-dimethoxy-17-hydroxy-7-methyl-19-norpregn-5(10)-en-20-yn-3-one (15 kg) in a mixture of pyridine (630 ml) and ethanol (315 litres) was mixed with a solution of oxalic acid (750 gr) in water (90 litres) and the mixture was stirred for 2 hours at approximately 30 °C. The solution was poured out in a mixture of pyridine (1350 ml) and water (300 litres) and the resulting suspension was filtered. The crystals were washed with a mixture of water and pyridine and dried under vacuum at 40 °C to give $(7\alpha,17\alpha)$ -17-hydroxy-7-methyl-19-norpregn-5(10)-en-20-yn-3-one containing 0.6% of the corresponding $(7\alpha,17\alpha)$ -17-hydroxy-7-methyl-19-norpregn-4-en-20-yn-3-one as indicated by HPLC analysis; a stress test at 45 °C (duration 1 month) indicated a 0.4% increase of the latter compound.

Example 2

A solution of $(7\alpha,17\alpha)$ -3,3-dimethoxy-17-hydroxy-7-methyl-19-norpregn-5(10)-en-20-yn-3-one (15 kg) in a mixture of pyridine (630 ml) and ethanol (315 litres) was mixed with a solution of oxalic acid (375 gr) in water (90 litres) and the mixture was stirred for 3 hours at approximately 30 °C. The solution was poured out in a mixture of pyridine (1350 ml) and water (300 litres) and the resulting suspension is filtered. The crystals are washed with a mixture of water and pyridine and allowed to age for 3-6 days at room temperature. Subsequently, the crystals were dried under vacuum at 40 °C to give $(7\alpha,17\alpha)$ -17-hydroxy-7-methyl-19-norpregn-5(10)-en-20-yn-3-one containing \leq 0.1% of the corresponding $(7\alpha,17\alpha)$ -17-hydroxy-7-methyl-19-norpregn-4-en-20-yn-3-one as indicated by HPLC analysis; a stress test at 45 °C (duration 1 week) indicated a < 0.1% increase of the latter compound.

Example 3

The preparation as described in example 2 was repeated. $(7\alpha,17\alpha)$ -17-hydroxy-7-methyl-19-norpregn-5(10)-en-20-yn-3-one was obtained which contained 0.2 % of the corresponding $(7\alpha,17\alpha)$ -17-hydroxy-7-methyl-19-norpregn-4-en-20-yn-3-one as indicated by HPLC analysis; a stress test at 45 °C (duration 1 week) indicated a 0.1% increase of the latter compound.

Example 4

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A basic granulate was prepared by granulation of a mixture of lactose (diluent), potato starch (disintegrant) and potato starch mucilage (binder) in a fluid bed granulator. The water content of the granulate varied within 5.5% - 6.5%. After granulation, the basic granulate was passed through a conical high speed sieve. Part of the granulate (10% w/w) was mixed with Tibolone and ascorbyl palmitate using a tumble blender and then passed through a conical high speed sieve.

The Tibolone premix and the remainder of the basic granulate were mixed in a ribbon blender. Magnesium stearate was added and mixed. The final granulate was compressed into round tablets.

The stability of the active compound (Tibolone) in tablets was determined.

Table 1: Content of decomposition product (Org OM38) in percentage of the declared amount of Tibolone per tablet, in tablets containing a various amount of Tibolone, after storage at 25°C and 60% relative humidity.

	Concentration of Tibolone per tablet										
	0.46	2.5									
Storage time	Amount of Org OM38 formed during storage (in percentage of the										
(months)		declared amount of tibolone)									
0	1.2	0.8	. 0.5.	0.4							
6	6.5	3.5	1.8	1.6							
12	9.5	5.1	2:7	2.2							
18	12.2	6.1	3.3	2.7							

Example 5

Tablets of 1.25 mg of Tibolone have been prepared as described in example 4. The tablets were stored at 25°C and 60% relative humidity and the decomposition product (Org OM38) was measured.

Table 2: Content of decomposition product (Org OM38) in percentage of the declared amount of Tibolone per tablet. Stability of three development tablet batches (1.25 mg of Tibolone per 65 mg) was assessed (storage at 25°C and 60% relative humidity).

		Batch no									
	049514001	049515001	049516001								
Storage time		38 formed during storag									
(months)	de	declared amount of Tibolone)									
0	0.7;	Ϊ.0	1.3								
6	2.3	2.6	2.9								
12	3.5	3.7	3.8								
18	4.3	4.2	4.3								
. 24	5.1	4.9	4.9								

It can be concluded that the shelf life of tablets containing 1.25 mg of Tibolone per tablet of 65 mg is borderline.

Example 6

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Tibolone as prepared as in example 2 was used as the active compound to prepare tablets as described in example 4. The amount of Org OM38 formed in several batches during storage was determined.

Table 3: The stability of six tablet batches (1.25 mg of Tibolone per 65 mg) was assessed (storage at 25°C and 60% relative humidity). The amount of water incorporated in the basic granulate was varied from 6.0% to 6.5%.

			Batc	h no								
	TD96.1128	TD96.1132	TD96.1133	162454001	162455001	162456001						
Storage time	Amount	of Org OM	88 formed du	ring storage	(in percenta	ge of the						
(months)	·	declared amount of Tibolone)										
0	0.7	0.5	0.5	0.9	0.8	0.9						
6	1.3	1.1	1.1	1.8	1.7	1.8						
12	1.8	1.5	1.6									
18	2.0	1.5	1.7									
Water	6.5	6.5	6.5	6.3	6.1	6.1						
content of		}	ļ.									
the basic						1						
granulate					•							

Claims

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- 1. A high purity composition comprising (7α,17α)-17-hydroxy-7-methyl-19-nor-17-pregn-5(10)-en-20-yn-3-one, characterized in that the said composition comprises (7α,17α)-17-hydroxy-7-methyl-19-nor-17-pregn-4-en-20-yn-3-one in an amount less than 0.5%.
- The composition according to claim 1 characterized in that the amount of (7α,17α) 17-hydroxy-7-methyl-19-nor-17-pregn-4-en-20-yn-3-one is 0.25% or less.
- 3. The composition according to claim 1 characterized in that the amount of $(7\alpha,17\alpha)$ 17-hydroxy-7-methyl-19-nor-17-pregn-4-en-20-yn-3-one is 0.1% or less.
 - 4. A process for preparing the high purity compositions of claims 1-3 characterized in that crystals of (7α,17α)-17-hydroxy-7-methyl-19-nor-17-pregn-5(10)-en-20-yn-3one are allowed to age in the presence of water for at least 24 hours.
 - 5. The process according to claim 4 wherein the aging lasts 3-6 days.
- 15 6. The process according to claims 4 or 5 characterized in that the crystals are formed in the last step of the Tibolone synthesis comprising the steps of
 - a. reacting $(7\alpha,17\alpha)$ -3,3-dimethoxy-17-hydroxy-7-methyl-19-norpregn-5(10)-en-20-yn-3-one in an organic solvent with a weak acidic aqueous solution b. pouring out the solution in water which is made slightly alkaline
 - c. washing the crystals with water which is made slightly alkaline.
 - 7. A pharmaceutical dosage unit obtainable by admixture of a pharmaceutically suitable solid carrier and the composition according to any one of the claims 1-3.
 - 8. A pharmaceutical dosage unit obtainable by admixture of a pharmaceutically suitable solid carrier and the composition obtainable by the process of claims 4-6.
- A dosage unit comprising a pharmaceutically suitable solid carrier and (7α,17α)-17-hydroxy-7-methyl-19-nor-17-pregn-5(10)-en-20-yn-3-one in an amount of less than
 2.50 mg and having a shelf life specification comprising less than 5% of (7α,17α)-17-hydroxy-7-methyl-19-nor-17-pregn-4-en-20-yn-3-one.
- 10. The dosage unit according to claim 9 characterized in that (7α,17α)-17-hydroxy-7 30 methyl-19-nor-17-pregn-5(10)-en-20-yn-3-one is present in an amount of 1.25 mg or less.

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- 11. The dosage unit according to claim 9 characterized in that $(7\alpha,17\alpha)$ -17-hydroxy-7-methyl-19-nor-17-pregn-5(10)-en-20-yn-3-one is present in an amount of 0.625 mg or less.
- 12. The dosage unit according to claims 9-11 wherein the shelf life is 1.5, more preferably 2 years.
- 13. The dosage unit according to claim 9-11 wherein at a shelf life period of 6 months the amount of (7α,17α)-17-hydroxy-7-methyl-19-nor-17-pregn-4-en-20-yn-3-one is 3 % or less, more preferably 2% or less.
- 14. The dosage unit according to claim 13 wherein the shelf life period is 1, preferably 1 ½ year, more preferably 2 years.

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	ENTS CONSIDERED TO BE RELEVANT	· · · · · · · · · · · · · · · · · · ·		
Category ?	Citation of document, with indication, where appropriate, of the rel	evant passages		Relevant to daim No.
X	EP 0 389 035 A (AKZO NV)		1	1-3,7-9,
	26 September 1990 (1990-09-26) examples 3,5,8	4jt		12-14
X	WO 98 47517 A (MORITA RYOICHI ;AK			9-14
	NV (NL); HAAN DE PIETER (NL); LAN	MBREGTS)		
	29 October 1998 (1998-10-29) page 2, last paragraph; examples	1-4 6-11		
		1 4,0 11		
X	WO 98 39012 A (AKZO NOBEL NV ;ZAN		·	9-14
	PIETER (NL); MEULEMAN DIRK GERRIT	「 (NL))		
	ll September 1998 (1998-09-11) page 11; example 1; table II			
	page 11, example 1, table 11			
χ	WO 89 09058 A (AKZO NV)		-	9
	5 October 1989 (1989-10-05)			
`	example 1			
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	an the priority date claimed	"&" document member	of the same patent f	amily
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EP 0 707 848 A (AKZO NOBEL NV) 24 April 1996 (1996-04-24) example 3 EP 0 613 687 A (AKZO NOBEL NV) 7 September 1994 (1994-09-07) example 2 EP 0 159 739 A (AKZO NV) 30 October 1985 (1985-10-30) example 1 A VAN VLIET N P ET AL: "An alternative synthesis of 17. betahydroxy-7.alphamethyl-19-nor-17.alphapregn-5(10)-en-20-yn-3-one (Org 0D 14)" RECUEIL DES TRAVAUX CHIMIQUES DES PAYS-8AS., vol. 105, no. 4, April 1986 (1986-04), pages 111-115, XP002099865 AMSTERDAM NL page 113, column 1, last paragraph page 115, column 1, paragraph 4 DECLERCO J P ET AL: "Conformational analysis of 3-oxo 5(10)-unsaturated steroids. Single-crystal x-ray structure analysis of 17-hydroxy-7.alphamethyl-19-nor-17.alphapregn-5(10)-en-20-yn-3-o ne (Org 0D 14)" RECUEIL DES TRAVAUX CHIMIQUES DES PAYS-8AS., vol. 103, no. 5, May 1984 (1984-05), pages 145-147, XP002099866 AMSTERDAM NL page 146, column 1; table I A WIELAND P ET AL: "Steroids. CCXI. Synthesis of 7.alphamethyl-3-oxo-19-norandrosta -4, 9, 11-trienes" HELVETICA CHIMICA ACTA, vol. 50, no. 6, 21 September 1967 (1967-09-21), pages 1453-1461, XP002099867 BASEL CH page 1459, paragraph 1													_ F	CT/E	P 99	0/07	768		
EP 0 707 848 A (AKZO NOBEL NV) 24 April 1996 (1996-04-24) example 3 EP 0 613 687 A (AKZO NOBEL NV) 7 September 1994 (1994-09-07) example 2 EP 0 159 739 A (AKZO NV) 30 October 1985 (1985-10-30) example 1 A VAN VLIET N P ET AL: "An alternative synthesis of 17.betahydroxy-7.alphamethyl-19-nor -17.alphapregn-5(10)-en-20-yn-3-one (Org 0D 14)" RECUEIL DES TRAVAUX CHIMIQUES DES PAYS-BAS., vol. 105, no. 4, April 1986 (1986-04), pages 111-115, XP002099865 AMSTERDAM NL page 113, column 1, last paragraph page 115, column 1, paragraph 4 A DECLERCO J P ET AL: "Conformational analysis of 3-oxo 5(10)-unsaturated steroids. Single-crystal x-ray structure analysis of 17-hydroxy-7.alphamethyl -19-nor-17.alphapregn-5(10)-en-20-yn-3-o ne (Org 0D 14)" RECUEIL DES TRAVAUX CHIMIQUES DES PAYS-BAS., vol. 103, no. 5, May 1984 (1984-05), pages 145-147, XP002099866 AMSTERDAM NL page 146, column 1; table I A MIELAND P ET AL: "Steroids. CCXI. Synthesis of 7.alphamethyl-3-oxo-19-norandrosta -4, 9, 11-trienes" HELVETICA CHIMICA ACTA, vol. 50, no. 6, 21 September 1967 (1967-09-21), pages 1453-1461, XP002099867 BASEL CH page 1459, paragraph 1 US 3 340 279 A (H. P. DE JONGH ET AL) 5 September 1967 (1967-09-05)	DOCUM	CUME	ENTS	CON	SIDERI	ED TO	BE RE	LEVA	NT										
24 April 1996 (1996-04-24) example 3 EP 0 613 687 A (AKZO NOBEL NV) 7 September 1994 (1994-09-07) example 2 EP 0 159 739 A (AKZO NV) 30 0ctober 1985 (1985-10-30) example 1 VAN VLIET N P ET AL: "An alternative synthesis of 17. betahydroxy-7.alphamethyl-19-nor -17.alphapregn-5(10)-en-20-yn-3-one (Org 0D 14)" RECUEIL DES TRAVAUX CHIMIQUES DES PAYS-8AS., vol. 105, no. 4, April 1986 (1986-04), pages 111-115, KP002099865 AMSTERDAM NL page 113, column 1, last paragraph page 115, column 1, paragraph 4 DECLERCO J P ET AL: "Conformational analysis of 3-oxo 5(10)-unsaturated steroids. Single-crystal x-ray structure analysis of 17-hydroxy-7.alphamethyl -19-nor-17.alphapregn-5(10)-en-20-yn-3-o ne (Org 0D 14)" RECUEIL DES TRAVAUX CHIMIQUES DES PAYS-8AS., vol. 103, no. 5, May 1984 (1984-05), pages 145-147, XP002099866 AMSTERDAM NL page 146, column 1; table I WIELAND P ET AL: "Steroids. CCXI. Synthesis of 7.alphamethyl-3-oxo-19-norandrosta -4, 9,11-trienes" HELVETICA CHIMICA ACTA, vol. 50, no. 6, 21 September 1967 (1967-09-21), pages 1453-1461, XP002099867 BASEL CH page 1459, paragraph 1 US 3 340 279 A (H. P. DE JONGH ET AL) 5 September 1967 (1967-09-05)	ition of do	of docu	cumer	nt, with	n indica	tion,wh	iere app	propria	te. of the	releva	ant pass	ages	•			Relev	ant to da	aim No.	
7 September 1994 (1994-09-07) example 2 EP 0 159 739 A (AKZO NV) 30 October 1985 (1985-10-30) example 1 VAN VLIET N P ET AL: "An alternative synthesis of 17.betahydroxy-7.alphamethyl-19-nor -17.alphapregn-5(10)-en-20-yn-3-one (Org -17.alphapregn-5(10)-en-20-yn-3-one (Org -17.alphapregn-5(10)-en-20-yn-3-one (Org -17.alphapregn-5(10)-en-20-yn-3-one (Org -18.column 1, paragraph -19.column 1, last paragraph -19.column 1, last paragraph -19.column 1, paragraph 4 DECLERCO J P ET AL: "Conformational analysis of 3-oxo 5(10)-unsaturated steroids. Single-crystal x-ray structure analysis of 17-hydroxy-7.alphamethyl -19-nor-17.alphapregn-5(10)-en-20-yn-3-o ne (Org OD 14)" RECUEIL DES TRAVAUX CHIMIOUES DES PAYS-BAS., vol. 103, no. 5, May 1984 (1984-05), pages 145-147. XPO02099866 AMSTERDAM NL page 146, column 1; table I MIELAND P ET AL: "Steroids. CCXI. Synthesis of 7.alphamethyl-3-oxo-19-norandrosta -4,9,11-trienes" HELVETICA CHIMICA ACTA, vol. 50, no. 6, 21 September 1967 (1967-09-21), pages 1453-1461, XPO02099867 BASEL CH page 1459, paragraph 1 US 3 340 279 A (H. P. DE JONGH ET AL) 5 September 1967 (1967-09-05)	24 Ap	Apr	ril	199													9-14		
30 October 1985 (1985-10-30) example 1 VAN VLIET N P ET AL: "An alternative synthesis of 17.betahydroxy-7.alphamethyl-19-nor -17.alphapregn-5(10)-en-20-yn-3-one (Org OD 14)" RECUEIL DES TRAVAUX CHIMIQUES DES PAYS-BAS., vol. 105, no. 4, April 1986 (1986-04), pages 111-115, XP002099865 AMSTERDAM NL page 113, column 1, last paragraph page 115, column 1, paragraph 4 DECLERCO J P ET AL: "Conformational analysis of 3-oxo 5(10)-unsaturated steroids. Single-crystal x-ray structure analysis of 17-hydroxy-7.alphamethyl -19-nor-17.alphapregn-5(10)-en-20-yn-3-o ne (Org OD 14)" RECUEIL DES TRAVAUX CHIMIQUES DES PAYS-BAS., vol. 103, no. 5, May 1984 (1984-05), pages 145-147, XP002099866 AMSTERDAM NL page 146, column 1; table I MIELAND P ET AL: "Steroids. CCXI. Synthesis of 7.alphamethyl-3-oxo-19-norandrosta -4,9,11-trienes" HELVETICA CHIMICA ACTA, vol. 50, no. 6, 21 September 1967 (1967-09-21), pages 1453-1461, XP002099867 BASEL CH page 1459, paragraph 1 US 3 340 279 A (H. P. DE JONGH ET AL) 5 September 1967 (1967-09-05)	7 Sep	Sept	teml	ber											•		9		
synthesis of 17.betahydroxy-7.alphamethyl-19-nor -17.alphapregn-5(10)-en-20-yn-3-one (Org 0D 14)" RECUEIL DES TRAVAUX CHIMIQUES DES PAYS-BAS., vol. 105, no. 4, April 1986 (1986-04), pages 111-115, XP002099865 AMSTERDAM NL page 113, column 1, last paragraph page 115, column 1, paragraph 4 —— DECLERCO J P ET AL: "Conformational analysis of 3-oxo 5(10)-unsaturated steroids. Single-crystal x-ray structure analysis of 17-hydroxy-7.alphamethyl -19-nor-17.alphapregn-5(10)-en-20-yn-3-o ne (Org OD 14)" RECUEIL DES TRAVAUX CHIMIQUES DES PAYS-BAS., vol. 103, no. 5, May 1984 (1984-05), pages 145-147, XP002099866 AMSTERDAM NL page 146, column 1; table I —— WIELAND P ET AL: "Steroids. CCXI. Synthesis of 7.alphamethyl-3-oxo-19-norandrosta -4,9,11-trienes" HELVETICA CHIMICA ACTA, vol. 50, no. 6, 21 September 1967 (1967-09-21), pages 1453-1461, XP002099867 BASEL CH page 1459, paragraph 1 US 3 340 279 A (H. P. DE JONGH ET AL) 5 September 1967 (1967-09-05)	30 Oc	0ct	tobe	er :				•	10)		•	. •					9		
analysis of 3-oxo 5(10)-unsaturated steroids. Single-crystal x-ray structure analysis of 17-hydroxy-7.alphamethyl -19-nor-17.alphapregn-5(10)-en-20-yn-3-o ne (Org OD 14)" RECUEIL DES TRAVAUX CHIMIQUES DES PAYS-BAS., vol. 103, no. 5, May 1984 (1984-05), pages 145-147, XPO02099866 AMSTERDAM NL page 146, column 1; table I WIELAND P ET AL: "Steroids. CCXI. Synthesis of 7.alphamethyl-3-oxo-19-norandrosta -4,9,11-trienes" HELVETICA CHIMICA ACTA, vol. 50, no. 6, 21 September 1967 (1967-09-21), pages 1453-1461, XPO02099867 BASEL CH page 1459, paragraph 1 US 3 340 279 A (H. P. DE JONGH ET AL) 5 September 1967 (1967-09-05)	synth 17.be -17.a OD 14 RECUE PAYS- vol. pages AMSTE page	nthe .bet 7.al 14) CUEI YS-B 1. 1 ges STER	esis ta lpha)" IL [BAS. 105; 1113	s of hydap DES , nd 1-11	f drox oregi TRA O. 4 O. 4 NL Olumi	y-7. n-5(VAU) , Ar XPO(alph (10)- (CH: pril)2099	ha -en- IMIQ 198 9865 st p	meth 20-y UES 6 (1	y1-1 n-3- DES 986- raph	19-no -one -04),	(0rg		. "	į		1-14		
Synthesis of 7.alphamethyl-3-oxo-19-norandrosta -4,9,11-trienes" HELVETICA CHIMICA ACTA, vol. 50, no. 6, 21 September 1967 (1967-09-21), pages 1453-1461, XP002099867 BASEL CH page 1459, paragraph 1 US 3 340 279 A (H. P. DE JONGH ET AL) 5 September 1967 (1967-09-05)	analy stero analy -19-n ne (0 RECUE PAYS- vol. 145-1	alys eroi alys 9-no (Or CUEI YS-B 1. 1 5-14 STER	sis ids. sis or-1 rg (IL [BAS. 103, RDAN	of of 17.a DD 1 DES , no XPO	3-02 ingle 17-1 alpha 14)" TRAV 0. 5 00209 NL	xo 5 e-cr hydr a -p VAUX , Ma 9986	icysta roxy- pregr CHI)-un al x -7.a n-5(IMIQ 984	ray Ipha 10)- UES	rate str me en-2 DES	ed ructu ethyl 20-yr	-3-0					1-14		
5 September 1967 (1967-09-05)	Synth 7.alp -4,9, HELVE vol. 21 Se 1453- BASEL	nthe alph ,9,1 LVET I. 5 Sep 53-1 SEL	esis ha 11-t TIC/ 50, oten 1461 CH	s of met trie A Ch no. nber l,)	thyl- thyl- enes' HIMI(6, 196 (P00)	-3-0 " CA A 57 (2099	0x0-1 ACTA 1967 0867	19-n ,	oran	dros	sta						1-14		
· -/	5. Sep	Sept	temt	oer	1967					ET -/	'AL)					-	1-14		

Int. Ational Application No
PCT/EP 99/07768

		PCT/EP 99/07768
C.(Contin Category	uation) DOCUMENTS CONSIDERED TO BE RELEVANT Citation of document, with indication where appropriate, of the relevant passages	lo.
	appropriate. of the felevant passages	Relevant to claim No.
A .	FR 1 583 441 A (THE UPJOHN COMPANY) 31 October 1969 (1969-10-31) example 16	1-14
Α .	US 3 475 465 A (WINTER MAX SOLOMON DE ET AL) 28 October 1969 (1969-10-28) example II	1-14
4 ·	US 3 432 528 A (ANNER GEORG ET AL) 11 March 1969 (1969-03-11) example 3	1-14
A	US 3 576 828 A (ANNER GEORG ET AL) 27 April 1971 (1971-04-27) example 9	1-14
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Information on patent family members

Inte ional Application No PCT/EP 99/07768

				000	73, 0, , 00
Patent document cited in search report		Publication date		Patent family member(s)	Publication date
<u>'</u>		·			<u> </u>
EP 0389035	Α	26-09-1990	AT	98490 T	. 15-01-1994
		•	AU	625083 B	02-07-1992
			AU	5139490 A	20-09-1990
			CA	2011452 A	18-09-1990
			DE	69005165 D	
					27-01-1994
			DE	69005165 T	05-05-1994
			ÐK	389035 T	14-02-1994
		•	ES	2062292 T	16-12-1994
			FI	94865 B	31-07-1995
			HK	1002019 A	24-07-1998
			IE	63051 B	22-03-1995
			JP	3047195 A	
					28-02-1991
		•	. MX	9203787 A	01-07-1992
			NZ	232946 A	27-08-1991
		•	PT	93488 A,B	07-11-1990
•			US	5037817 A	06-08-1991
WO 9847517	Λ	20-10-1000		0014600 4	10.11.100
MU 304/31/	Α	29-10-1998	AU	8014698 A	13-11-1998
			JP	10316573 A	02-12-1998
•			NO	995127 A	21-10-1999
			ZA	9803169 A	20-10-1998
WO 9839012	Α	11-09-1998	AU	6720400 4	22 00 1000
## JOJJU16 .	Λ	11 03-1330		6728498 .A	22-09-1998
			. ZA 	9801731 A	07-09-1998
WO 8909058	Α	05-10-1989	AU	3435989 A	16-10-1989
			DK	224690 A	18-09-1990
			EP	0406279 A	09-01-1991
			JP	3503414 T	01-08-1991
ED 0707040		24 04 1005			
EP 0707848	A	24-04-1996	AU	688581 B	12-03-1998
	٠.		· AU	3426795 A	02-05-1996
			BR	9504400 A	27-05-1997
			CA	2159419 A	18-04-1996
			CN	1130064 A	04-09-1996
			FI	954905 A	
					18-04-1996
			HU	75247 A	28-05-1997
		•	IL	115445 A	17-08-1999
			JP	8268914 A	15-10-1996
			TR.	960302 A	21-06-1996
EP 0613687	Α	07-09-1994	AT	180669 T	15.06.1000
-LI 001300/	n	01-03-1334			15-06-1999
			AU	671706 B	05-09-1996
			AU	5754294 A	08-09-1994
			CA	2116829 A	06-09-1994
			DE	69418744 D	08-07-1999
			DE	69418744 T	11-11-1999
	•		ES	2134313 T	01-10-1999
			JP	7002673 A	
		•			06-01-1995
			NO	940777 A	06-09-1994
			NZ	260017 A	24-06-1997
			. US	5512556 A	30-04-1996
			ZA	9401464 A	27-09-1994
ED 0150720	Δ	20 10 1005		4000 = =	
EP 0159739	A	30-10-1985	AT	42895 T	15-05-1989
			. JP	1829698 C	15-03-1994
			JP	60209599 A	22-10-1985
			MX	9203811 A	01-07-1992
	~				

Information on patent family members

Inte .ional Application No
PCT/EP 99/07768

				1		P 99/07768
	itent document I in search repoi	rt	Publication date		Patent family member(s)	Publication date
EP	0159739	Α		US	4701450 A	20-10-1987
US	3340279	'A	05-09-1967	NL BE	6406797 A 665514 A	17-12-1965
				CH	517725 A	15-01-1972
				DE	1543273 A	07-08-1969
		•		DK	114553 B	14-07-1969
				FR	1594513 A	08-06-1970
				GB	1104462 A	
				SE	336788 B	19-07-1971
FR	1583441	Α	31-10-1969	BE	712229 A	16-09-1968
				. CH.	516545 A	15-12-1971
				. DE	1668818 A	30-09-1971
				GB GB	1206872 A 1206873 A	30-09-1970
				GB	1206873 A 1206874 A	30-09-1970
	•			NL	6803328 A	30-09-1970 16-09-1968
				US	3515719 A	02-06-1970
US	3475465	Α.	28-10-1969	NL	6608779 A	27-12-1967
		-	}	BE	[₹] 700390 A	27-12-1967
	•	•		CH	537913.A.	31-07-1973
				DE	1618747 A	25-02-1971
				DK	115989 B	
				FR GB	1527563 A 1177845 A	06-11-1968
					11//049 M	14-01-1970
US	3432528	Α	11-03-1969	BE	684851 A	30-01-1967
				BE	684852 A	30-01-1967
				CH CH	488682 A	15-04-1970
				CH	509996 A 519488 A	15-07-1971 29-02-1972
				CH	525873 A	31-07-1972
	•			DE	1568308 A	05-02-1970
				DE	1568306 A	12-03-1970
				DE	1568307 A	05-02-1970
				FR	5260 M	24-07-1967
		•		FR	6350 M	07-10-1968
				FR FR	1491586 A 1491587 A	30-11-1967
				FR FR	1491587 A 1491588 A	30-11-1967 30-11-1967
				GB	1158331 A	16-07-1969
				GB	1158331 A	16-07-1969
				NL	6610741 A,B	31-01-1967
				NL.	6610742 A,B	31-01-1967
				NL	6610743 A.B	31-01-1967
				NL	7607401 A	29-10-1976
		•		US	3576828 A	27-04-1971
				ES.	329609 A	01-03-1968
				CS 	153439 B	25-02-1974
US :	3576828	Α	27-04-1971	BE	684851 A	30-01-1967
٠	•			BE	684852 A	30-01-1967
				CH	488682 A	15-04-1970
				CH CH	509996 A 519488 A	15-07-1971
				CH	525873 A	29-02-1972
				UII	3430/3 H	31-07-1972

Information on patent family members

Inte. Jonal Application No PCT/EP 99/07768

Patent document cited in search report	Publication date		t family ber(s)	Publication date
US 3576828 A		DE 1	568308 A	05-02-1970
		DE 1	568306 A	12-03-1970
		DE 1	568307 A	05-02-1970
·		· FR	5260 M	24-07-1967
		FR	6350 M	07-10-1968
·		FR 1	491586 A	30-11-1967
		FR 1	491587 A	30-11-1967
		FR 1	491588 A	30-11-1967
·		GB 1	158331 A	16-07-1969
•		GB 1	158332 A	16-07-1969
•		NL · 6	610741 A,B	31-01-1967
•		NL 6	610742 A,B	31-01-1967
:		. NL 6	610743 A,B	31-01-1967
		NL 7	607401 A	29-10-1976
		US 3	432528 A	11-03-1969
		ES	329609 A	01-03-1968
		CS	153439 B	25-02-1974

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